RAW SEQUENCE LISTING

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) no errors detected.

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IFWO

RAW SEQUENCE LISTING DATE: 02/22/2005
PATENT APPLICATION: US/10/803,580B TIME: 15:41:09

Input Set : A:\SYR-HDAC-5005-C2 sub seq list 2.ST25.txt

Output Set: N:\CRF4\02222005\J803580B.raw

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3 <110> APPLICANT: Syrrx, Inc.
     5 <120> TITLE OF INVENTION: HISTONE DEACETYLASE INHIBITORS
     7 <130> FILE REFERENCE: SYR-HDAC-5005-C2
     9 <140> CURRENT APPLICATION NUMBER: US 10/803,580B
    10 <141> CURRENT FILING DATE: 2004-03-17
    12 <150> PRIOR APPLICATION NUMBER: US 60/455,437
    13 <151> PRIOR FILING DATE: 2003-03-17
    15 <150> PRIOR APPLICATION NUMBER: US 60/531,203
    16 <151> PRIOR FILING DATE: 2003-12-19
    18 <160> NUMBER OF SEQ ID NOS: 8
    20 <170> SOFTWARE: PatentIn version 3.2
    22 <210> SEQ ID NO: 1
    23 <211> LENGTH: 513
    24 <212> TYPE: PRT
    25 <213> ORGANISM: Artificial
    27 <220> FEATURE:
    28 <223> OTHER INFORMATION: Residues 1-482 of HDAC1 with a
"MSYYHHHHHHDYDIPTTENLYFQGAMEPGGS"
    29
            tag at the N-terminus
    31 <400> SEQUENCE: 1
    33 Met Ser Tyr Tyr His His His His His Asp Tyr Asp Ile Pro Thr
    37 Thr Glu Asn Leu Tyr Phe Gln Gly Ala Met Glu Pro Gly Gly Ser Met
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    41 Ala Gln Thr Gln Gly Thr Arg Arg Lys Val Cys Tyr Tyr Tyr Asp Gly
                                    40
    45 Asp Val Gly Asn Tyr Tyr Tyr Gly Gln Gly His Pro Met Lys Pro His
                                55
    49 Arg Ile Arg Met Thr His Asn Leu Leu Leu Asn Tyr Gly Leu Tyr Arg
                            70
    53 Lys Met Glu Ile Tyr Arg Pro His Lys Ala Asn Ala Glu Glu Met Thr
                        85
                                            90
    57 Lys Tyr His Ser Asp Asp Tyr Ile Lys Phe Leu Arg Ser Ile Arg Pro
                   100
                                        105
                                                            110
    61 Asp Asn Met Ser Glu Tyr Ser Lys Gln Met Gln Arg Phe Asn Val Gly
                                    120
    62
               115
    65 Glu Asp Cys Pro Val Phe Asp Gly Leu Phe Glu Phe Cys Gln Leu Ser
           130
    69 Thr Gly Gly Ser Val Ala Ser Ala Val Lys Leu Asn Lys Gln Gln Thr
                                                155
                            150
    73 Asp Ile Ala Val Asn Trp Ala Gly Gly Leu His His Ala Lys Lys Ser
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                       165
    77 Glu Ala Ser Gly Phe Cys Tyr Val Asn Asp Ile Val Leu Ala Ile Leu
    78
                   180
                                        185
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81 Glu Leu Leu Lys Tyr His Gln Arg Val Leu Tyr Ile Asp Ile Asp Ile 200 85 His His Gly Asp Gly Val Glu Glu Ala Phe Tyr Thr Thr Asp Arg Val 215 89 Met Thr Val Ser Phe His Lys Tyr Gly Glu Tyr Phe Pro Gly Thr Gly 230 93 Asp Leu Arg Asp Ile Gly Ala Gly Lys Gly Lys Tyr Tyr Ala Val Asn 250 245 97 Tyr Pro Leu Arg Asp Gly Ile Asp Asp Glu Ser Tyr Glu Ala Ile Phe 265 101 Lys Pro Val Met Ser Lys Val Met Glu Met Phe Gln Pro Ser Ala Val 275 280 105 Val Leu Gln Cys Gly Ser Asp Ser Leu Ser Gly Asp Arg Leu Gly Cys 295 109 Phe Asn Leu Thr Ile Lys Gly His Ala Lys Cys Val Glu Phe Val Lys 310 315 113 Ser Phe Asn Leu Pro Met Leu Met Leu Gly Gly Gly Tyr Thr Ile 330 325 117 Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr Ala Val Ala Leu Asp 345 121 Thr Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp Tyr Phe Glu Tyr Phe 360 125 Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser Asn Met Thr Asn Gln 375 380 129 Asn Thr Asn Glu Tyr Leu Glu Lys Ile Lys Gln Arg Leu Phe Glu Asn 395 390 133 Leu Arg Met Leu Pro His Ala Pro Gly Val Gln Met Gln Ala Ile Pro 405 410 137 Glu Asp Ala Ile Pro Glu Glu Ser Gly Asp Glu Asp Glu Asp Pro 420 425 430 141 Asp Lys Arg Ile Ser Ile Cys Ser Ser Asp Lys Arg Ile Ala Cys Glu 435 145 Glu Glu Phe Ser Asp Ser Glu Glu Glu Glu Gly Gly Arg Lys Asn 455 149 Ser Ser Asn Phe Lys Lys Ala Lys Arg Val Lys Thr Glu Asp Glu Lys 470 475 153 Glu Lys Asp Pro Glu Glu Lys Lys Glu Val Thr Glu Glu Lys Thr 490 485 157 Lys Glu Glu Lys Pro Glu Ala Lys Gly Val Lys Glu Glu Val Lys Leu 161 Ala 165 <210> SEQ ID NO: 2 166 <211> LENGTH: 1542 167 <212> TYPE: DNA 168 <213> ORGANISM: Artificial 170 <220> FEATURE: 171 <223> OTHER INFORMATION: DNA sequence encoding residues 1-482 of HDAC1 with a "MSYYHHHHHHDYDIPTTENLYFQGAMEPGGS" tag at the N-terminus 174 <400> SEQUENCE: 2

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    175 atgtcgtact accatcacca tcaccatcac gattacgata tcccaacgac cgaaaacctg
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    177 tattttcaqq qcqccatqqa acccqgqqqa tccatggcgc agacgcaggg cacccggagg
    179 aaagtetgtt actactacga eggggatgtt ggaaattact attatggaca aggecaceca
                                                                               180
    181 atquaqcete accquatecq catqueteat autttgetge teaactatgg tetetaccqu
    183 aaaatqqaaa tctatcgccc tcacaaagcc aatgctgagg agatgaccaa gtaccacagc
                                                                               300
                                                                               360
    185 gatgactaca ttaaattett gegeteeate egteeagata acatgtegga gtacageaag
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    187 cagatgcaga gattcaacgt tggtgaggac tgtccagtat tcgatggcct gtttgagttc
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    189 tgtcagttgt ctactggtgg ttctgtggca agtgctgtga aacttaataa gcagcagacg
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    191 qacatcqctg tgaattgggc tgggggcctg caccatgcaa agaagtccga ggcatctggc
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    195 gtgctgtaca ttgacattga tattcaccat ggtgacggcg tggaagaggc cttctacacc
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    197 acggaccggg tcatgactgt gtcctttcat aagtatggag agtacttccc aggaactggg
    199 gacctacggg atatcggggc tggcaaaggc aagtattatg ctgttaacta cccgctccga
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    201 gacgggattg atgacgagtc ctatgaggcc attttcaagc cggtcatgtc caaagtaatg
    203 gagatgttcc agcctagtgc ggtggtctta cagtgtggct cagactccct atctggggat
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    205 cggttaggtt gcttcaatct aactatcaaa ggacacgcca agtgtgtgga atttgtcaag
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    207 agetttaace tgeetatget gatgetggga ggeggtggtt acaccatteg taacgttgee
    209 cqqtqctqqa catatqaqac aqctqtggcc ctggatacgg agatccctaa tgagcttcca
                                                                              1080
                                                                              1140
    211 tacaatqact actttqaata ctttqqacca gatttcaagc tccacatcag tccttccaat
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    213 atgactaacc agaacacgaa tgagtacctg gagaagatca aacagcgact gtttgagaac
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    215 cttagaatgc tgccgcacgc acctggggtc caaatgcagg cgattcctga ggacgccatc
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    217 cctgaggaga gtggcgatga ggacgaagac gaccctgaca agcgcatctc gatctgctcc
    219 tctgacaaac gaattgcctg tgaggaagag ttctccgatt ctgaagagga gggagagggg
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    221 ggccgcaaga actcttccaa cttcaaaaaa gccaagagag tcaaaacaga ggatgaaaaa
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    223 gagaaagacc cagaggagaa gaaagaagtc accgaagagg agaaaaccaa ggaggagaag
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     225 ccagaagcca aaggggtcaa ggaggaggtc aagttggcct ga
     228 <210> SEQ ID NO: 3
     229 <211> LENGTH: 498
     230 <212> TYPE: PRT
     231 <213> ORGANISM: Artificial
     233 <220> FEATURE:
     234 <223> OTHER INFORMATION: Residues 1-488 of HDAC2 with a "GHHHHHHH" tag at the C-
terminus
               and a "MGS" tag at the N-terminus
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     237 <400> SEQUENCE: 3
     239 Met Gly Ser Met Ala Tyr Ser Gln Gly Gly Lys Lys Lys Val Cys
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     243 Tyr Tyr Tyr Asp Gly Asp Ile Gly Asn Tyr Tyr Tyr Gly Gln Gly His
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     247 Pro Met Lys Pro His Arg Ile Arg Met Thr His Asn Leu Leu Leu Asn
     248
                 35
                                     40
     251 Tyr Gly Leu Tyr Arg Lys Met Glu Ile Tyr Arg Pro His Lys Ala Thr
                                 55
     252
     255 Ala Glu Glu Met Thr Lys Tyr His Ser Asp Glu Tyr Ile Lys Phe Leu
                                                                      80
                             70
                                                 75
     259 Arg Ser Ile Arg Pro Asp Asn Met Ser Glu Tyr Ser Lys Gln Met Gln
                                             90
     260
     263 Arg Phe Asn Val Gly Glu Asp Cys Pro Val Phe Asp Gly Leu Phe Glu
                                         105
     264
     267 Phe Cys Gln Leu Ser Thr Gly Gly Ser Val Ala Gly Ala Val Lys Leu
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Output Set: N:\CRF4\02222005\J803580B.raw

115
130
275 His Ala Lys Lys Ser Glu Ala Ser Gly Phe Cys Tyr Val Asn Asp Ile 150 155 160 155 160 160 175 160 160 175 160 175 160 175 160 175 170 175 175 160 175 170 175 175 175 170 175 185 175
160 145
279 Val Leu Ala Ile Leu Glu Leu Leu Lys Tyr His Gln Arg Val Leu Tyr 280
280
283 Ile Asp Ile Asp Ile His His Gly Asp Gly Val Glu Glu Ala Phe Tyr 284 180 185 185 190 287 Thr Thr Asp Arg Val Met Thr Val Ser Phe His Lys Tyr Gly Glu Tyr 205 205 288 195 200 200 205 205 291 Phe Pro Gly Thr Gly Asp Leu Arg Asp Ile Gly Ala Gly Lys Gly Lys 220 210 215 220 220 295 Tyr Tyr Ala Val Asn Phe Pro Met Arg Asp Gly Ile Asp Asp Glu Ser 230 230 235 240 299 Tyr Gly Gln Ile Phe Lys Pro Ile Ile Ser Lys Val Met Glu Met Tyr 245 260 255 240 303 Gln Pro Ser Ala Val Val Leu Gln Cys Gly Ala Asp Ser Leu Ser Gly 255 260 265 270 304 260 260 265 265 270 304 275 280 275 280 311 Val Glu Val Val Lys Thr Phe Asn Leu Pro Leu Leu Met Leu Met Leu Gly Gly 295 285 311 Val Gly Tyr Thr Ile Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr 316 300 315 Gly Gly Tyr Thr Ile Arg Asn Val Ala Arg Cys Trp Thr Tyr Asn Asp 325
284
287 Thr Thr Asp Arg Val Met Thr Val Ser Phe His Lys Tyr Gly Glu Tyr 288
288 195 200 205 205 205 291 291 Thr Gly Asp Leu Arg Asp Ile Gly Ala Gly Lys Gly Lys 220 210 210 215 215 220 225 225 225 225 220 225 220 220 220 220 220 220 220 220 220 220 220 220 220 220 220 220 220
291 Phe Pro Gly Thr Gly Asp Leu Arg Asp Ile Gly Ala Gly Lys Gly Lys 292
292 210 215 220 220 280 295 380 Phe Pro Met Arg Asp Gly Ile Asp Asp Glu Ser 240 290 225 230 230 235 235 250 240 240 240 290 297 Tyr Gly Gly Gly Fle Leu Ser 240 235 235 240 250 250 250 250 250 250 250 250 250 250 250 260 260 260 260 260 260 270 270 270 260 270 280 280 280 280 280 280 280 2
295 Tyr Tyr Ala Val Asn Phe Pro Met Arg Asp Gly Ile Asp Asp Gly Ile Asp Asp Gly Ile Asp Asp Ile Asp Asp Ile Asp Ile Ile Ile Ser Lys Val Met Glu Met Tyr 300 Ile Ile Ile Ser Lys Val Met Glu Met Tyr 300 Ile Ile Ile Ile Ser Lys Val Met Ile Ile Ile Ile Ile Ile Ile Asp Ile
296 225
299 Tyr Gly Gln Ile Phe Lys Pro Ile Ile Ser Lys Val Met Glu Met Tyr 300
300
303 Gln Pro Ser Ala Val Val Leu Gln Cys Gly Ala Asp Ser Leu Ser Gly 304
304
307 Asp Arg Leu Gly Cys Phe Asn Leu Thr Val Lys Gly His Ala Lys Cys 308
308
311 Val Glu Val Val Lys Thr Phe Asn Leu Pro Leu Leu Met Leu Gly Gly 312 290 295 300 315 Gly Gly Tyr Thr Ile Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr 316 305 310 310 315 320 319 Ala Val Ala Leu Asp Cys Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp 320 325 330 335 323 Tyr Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 340 325 345 345
312 290 295 300 315 Gly Gly Tyr Thr Ile Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr 316 305 317 Ala Val Ala Leu Asp Cys Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp 320 325 325 325 345 Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 355 365 375 385 385 385 385 385 385
315 Gly Gly Tyr Thr Ile Arg Asn Val Ala Arg Cys Trp Thr Tyr Glu Thr 316 305
316 305 310 315 320 319 Ala Val Ala Leu Asp Cys Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp 320 325 330 335 323 Tyr Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 340 345 345
319 Ala Val Ala Leu Asp Cys Glu Ile Pro Asn Glu Leu Pro Tyr Asn Asp 320 325 330 335 323 Tyr Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 345 345 350
320 325 330 335 323 Tyr Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 345 350
323 Tyr Phe Glu Tyr Phe Gly Pro Asp Phe Lys Leu His Ile Ser Pro Ser 324 340 345 350
324 340 345 350
328 355 360 365
331 Arg Leu Phe Glu Asn Leu Arg Met Leu Pro His Ala Pro Gly Val Gln
332 370
335 Met Gln Ala Ile Pro Glu Asp Ala Val His Glu Asp Ser Gly Asp Glu
336 385 390 395 400
339 Asp Gly Glu Asp Pro Asp Lys Arg Ile Ser Ile Arg Ala Ser Asp Lys
340 405 410 415
343 Arg Ile Ala Cys Asp Glu Glu Phe Ser Asp Ser Glu Asp Glu Gly Glu
344 420 425 430
347 Gly Gly Arg Arg Asn Val Ala Asp His Lys Lys Gly Ala Lys Lys Ala
348 435 440 445
351 Arg Ile Glu Glu Asp Lys Lys Glu Thr Glu Asp Lys Lys Thr Asp Val
352 450 455 460
355 Lys Glu Glu Asp Lys Ser Lys Asp Asn Ser Gly Glu Lys Thr Asp Thr
356 465 470 475 480
359 Lys Gly Thr Lys Ser Glu Gln Leu Ser Asn Pro Gly His His His
360 485 490 495
363 His His
367 <210> SEQ ID NO: 4

DATE: 02/22/2005

TIME: 15:41:09

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Input Set : A:\SYR-HDAC-5005-C2 sub seq list 2.ST25.txt
                     Output Set: N:\CRF4\02222005\J803580B.raw
     368 <211> LENGTH: 1497
     369 <212> TYPE: DNA
     370 <213> ORGANISM: Artificial
     372 <220> FEATURE:
     373 <223> OTHER INFORMATION: DNA sequence encoding residues 1-488 of HDAC2 with a
"GHHHHHH"
               tag at the C-terminus and a "MGS" tag at the N-terminus
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     376 <400> SEQUENCE: 4
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     381 atgacccata acttgctgtt aaattatggc ttatacagaa aaatggaaat atataggccc
     383 cataaagcca ctgccgaaga aatgacaaaa tatcacagtg atgagtatat caaatttcta
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     385 cggtcaataa gaccagataa catgtctgag tatagtaagc agatgcagag atttaatgtt
     387 ggagaagatt gtccagtgtt tgatggactc tttgagtttt gtcagctctc aactggcggt
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     391 ggaggattac atcatgctaa gaaatcagaa gcatcaggat tctgttacgt taatgatatt
     393 qtqcttqcca tccttqaatt actaaagtat catcagagag tcttatatat tgatatagat
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     397 tcattccata aatatgggga atactttcct ggcacaggag acttgaggga tattggtgct
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     399 qqaaaaqqca aatactatqc tqtcaatttt ccaatgagag atggtataga tgatgagtca
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     401 tatqqqcaqa tatttaagcc tattatctca aaggtgatgg agatgtatca acctagtgct
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     403 qtqqtattac aqtqtqqtqc agactcatta tctgqtgata gactgggttg tttcaatcta
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     405 acaqtcaaaq qtcatqctaa atgtgtagaa gttgtaaaaa cttttaactt accattactg
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     409 gcagttgccc ttgattgtga gattcccaat gagttgccat ataatgatta ctttgagtat
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     411 tttqqaccaq acttcaaact qcatattagt ccttcaaaca tgacaaacca gaacactcca
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     413 qaatatatqq aaaaqataaa acagcgtttg tttgaaaatt tgcgcatgtt acctcatgca
     415 cctggtgtcc agatgcaagc tattccagaa gatgctgttc atgaagacag tggagatgaa
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     417 gatggagaag atccagacaa gagaatttct attcgagcat cagacaagcg gatagcttgt
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     419 gatgaagaat teteagatte tgaggatgaa ggagaaggag gtegaagaaa tgtggetgat
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     421 cataaqaaaq qaqcaaaqaa agctaqaatt gaagaagata agaaagaaac agaggacaaa
     423 aaaacagacg ttaaggaaga agataaatcc aaggacaaca gtggtgaaaa aacagatacc
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     425 aaaggaacca aatcagaaca gctcagcaac cccgggcatc accatcacca tcactaa
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     430 <212> TYPE: PRT
     431 <213> ORGANISM: Artificial
     433 <220> FEATURE:
     434 <223> OTHER INFORMATION: Residues 73-845 of HDAC6 with a "GHHHHHHH" tag at the C-
terminus
               and a "MP" tag at the N-terminus
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                     20
     447 Asp Ser Phe Pro Glu Gly Pro Glu Arg Leu His Ala Ile Lys Glu Gln
     448
                                     40
     451 Leu Ile Gln Glu Gly Leu Leu Asp Arg Cys Val Ser Phe Gln Ala Arg
     455 Phe Ala Glu Lys Glu Glu Leu Met Leu Val His Ser Leu Glu Tyr Ile
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RAW SEQUENCE LISTING

PATENT APPLICATION: US/10/803,580B

RAW SEQUENCE LISTING ERROR SUMMARY
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Invalid <213> Response:

Use of "Artificial" only as "<213> Organism" response is incomplete, per 1.823(b) of New Sequence Rules. Valid response is Artificial Sequence.

Seq#:1,2,3,4,5,6,7,8

VERIFICATION SUMMARY

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